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New Ar-Ar ages of southern Indian kimberlites and a lamproite and their geochemical evolution

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Abstract
The kimberlites and lamproites of southern India are thought to have formed in the most prolific known period of Precambrian ultramafic/ultrapotassic magmatism at around 1100 Ma. This study reports new age data for southern Indian ultrapotassic rocks (kimberlites and lamproites), a controversial topic due to the wide range of published age data and disagreements over the reliability of previously published ages. In this study we obtained new high-precision Ar-Ar data that better constrain the ages of southern Indian ultrapotassic rocks. Dates from three samples are presented, including two kimberlites from Wajrakarur kimberlite field and one lamproite from the Krishna lamproite field. These age data are then combined with bulk-rock geochemical and Nd isotopic data to provide further constraints on the source region and primary magma composition of southern Indian kimberlites and lamproites. Previously, the Chelima lamproite (ca. 1400 Ma) was considered to be one of the oldest lamproites in the world. However, our age data suggest that at least one lamproite (Pochampalle) was generated in the same region 100 Ma before the Chelima event. The Pochampalle lamproite was emplaced around ~1500 Ma as shown by the Ar-Ar data in this study, roughly 250 Ma before the other Krishna lamproites. It would seem that the Pochampalle lamproite was also derived from an isotopically distinct source region with a lower $^{143}\text{Nd}/^{144}\text{Nd}$ ratio than other lamproites in the Krishna field. These findings not only have implications for regional ultramafic/ultrapotassic magmatism, but also demonstrate that the mantle processes for producing lamproitic melts existed earlier than previously thought.

Keywords: Kimberlite, Lamproite, Eastern Dharwar Craton, Southern India, Proterozoic, Ar-Ar dating

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1. Introduction

Kimberlites and lamproites represent ultramafic magmas that travel rapidly from source regions in the deep mantle (>100 km) to the shallowest crustal levels at speeds of up to 0.1 to 4.0 m s\(^{-1}\) (Kelley & Wartho, 2000). In effect, they provide snapshots of the geochemical and isotopic nature of the deep lithospheric or asthenospheric mantle at the time of their emplacement. Because their petrogenesis has critical implications for large-scale processes (e.g. subduction, rifting, mantle plumes, lithospheric enrichment), accurate determination of their ages is important in order to investigate the evolution of the deep mantle through time.

The majority of known kimberlite and lamproite occurrences (>80%) globally are Mesozoic or Cenozoic in age and are often associated with periods of continental rifting and formation of ocean basins. For example, Cretaceous kimberlite and related magmatism in Southern Africa (125-55 Ma) closely follows the opening of the South Atlantic Ocean, which initiated around 130 Ma ago; similarly the majority of kimberlites and related ultramafic lamprophyres in Labrador, Quebec, West Greenland and Scandinavia were emplaced 630-560 Ma ago and linked to the Eocambrian opening of the Iapetus Ocean (Kumar et al., 2007). Heaman et al. (2003) indicate a lack of worldwide known kimberlite occurrences between 250 and 360 Ma that may be linked to relative crustal and mantle stability during the lifetime of the Gondwanaland supercontinent. It could therefore be proposed that a similar lack of kimberlite and lamproite occurrences of Proterozoic age is also due to crustal and mantle stability.
Proterozoic kimberlite and lamproites are relatively rare by comparison with Phanerozoic occurrences. The kimberlites and lamproites of southern India are thought to have formed in the most prolific known period of Precambrian ultramafic magmatism at around 1100 Ma. Kimberlites, lamproites and related alkalic rocks with emplacement ages of ~1100-1200 Ma have been reported from India, Canada, Greenland, Australia, Liberia, N. America, Scandinavia and South Africa (Kumar et al., 2007). Consequently this period of magmatism has been suggested to represent a global ‘mantle event’, when mantle conditions must have been favourable for kimberlite and lamproite emplacement (Skinner et al., 1985).

The kimberlites and lamproites of India have been proposed to be related to the formation of the Rodinia supercontinent (Kumar et al., 2007), which existed between 1000 and 750 Ma ago (Dalziel et al., 2000). The supercontinent began to form at ~1 Ga by accretion and collision of fragments produced by breakup of the older supercontinent, Columbia, which was assembled by global-scale 2.0-1.8 Ga collisional events (Zhao et al., 2004). Rodinia is thought to have included India, Australia and many of the present day continents of the southern hemisphere, which subsequently reorganized to form Gondwana.

The break-up of Columbia and initial assembly of Rodinia occurred ~1100 Ma ago, which may have provided favourable conditions for generation and emplacement of kimberlitic magmas across the Rodinian continent, producing many of the known Proterozoic kimberlites (~1100 Ma). During the formation of Rodinia ~1100 Ma ago, intracontinental rifting was taking place, resulting in the formation of several Large Igneous Provinces; e.g., 1109-1086 Ma in Laurentia (Ernst and Buchan, 2001)
and 1112-1102 Ma in the Kalahari Craton (Hanson et al., 2004). Kumar et al. (2007) also proposed an enhanced period of mantle plume activity, approximately 100 Ma prior to the formation of Rodinia.

The petrogenesis of the southern Indian kimberlites is currently still debated and several theories have been put forward. Chalapathi Rao et al. (2004) suggested that they are the result of lithospheric extension during the Mid-Proterozoic. This model proposes a sub-continental lithospheric source that experienced metasomatic enrichment caused by the migration of volatile-rich small fraction melts from the convecting mantle. The model also indicates that the kimberlites are the products of small-degree partial melting of a garnet peridotite mantle source, extensively depleted prior to metasomatic enrichment and subsequent partial melting (Chalapathi Rao et al., 2004). In contrast, Paton et al. (2009) suggested that the southern Indian kimberlites were derived directly from asthenospheric depths. Based on perovskite Sr and bulk-rock Hf and Nd isotopic data, these authors proposed that the asthenospheric component must have originated from within or below the transition zone, and may represent ancient subducted oceanic crust. Kumar et al. (2007) proposed that the southern Indian kimberlites located in the Eastern Dharwar craton (EDC) are likely to be linked to the impingement of a short-lived mantle plume beneath the Dharwar Craton, or that a major change and re-organization of the mantle convection regime occurred at the time of their formation.

There is also debate over whether lamproites are derived from the lithospheric (Davies et al., 2006) or convecting mantle (Murphy et al., 2002). Their complex geochemistry suggests a mantle source region that experienced both depletion and
enrichment. For the Krishna lamproites there are currently three main theories put forward for their formation. Chalapathi Rao et al. (2004) suggested that the source region underwent an extensive initial depletion involving komatiitic type melt extraction, followed by subsequent enrichment by a depleted MORB-type source. Paul et al. (2007) proposed that they were derived from partial melting of a mantle source that was metasomatically enriched in Ti and Fe. This same study also suggested that the lamproites were derived from depths shallower than the garnet stability field and their source regions were shallower than those of the kimberlites in the Eastern Dharwar Craton (EDC). Chakrabarti et al. (2007) maintained that the Krishna lamproites were derived by partial melting of metasomatized subducted Achaean komatiite, in a peridotite mantle-source assemblage. In their study, Chakrabarti et al. (2007) argued that the Pb-isotopic signatures (high $^{207}\text{Pb}$ low $^{206}\text{Pb}$) and superchondritic Nb/Ta ratios in Krishna lamproites ruled out their derivation from a metasomatized sub-continental lithospheric mantle source and instead suggested an Archean crustal component in the source (Chakrabarti et al., 2007).

There are relatively few known kimberlite and lamproite occurrences before ~1100 Ma, e.g., lamproite in Karelia, Russia 2.74 Ga (Sergreev et al., 2007); Kimberlite located in the Guyana Craton, Venezuela age dated at 1732 ± 82 using Rb-Sr chronometry (Nixon et al., 1992). The Proterozoic kimberlites and lamproites of southern India (Fig. 1) therefore provide a rare opportunity to study some of the oldest known occurrences of these rock types.

Previously published age data for the southern Indian kimberlites has been controversial, with different dating techniques showing wide ranges in age, while
several individual pipes remain undated. It has been proposed that the kimberlites in
the EDC had a contemporaneous emplacement at around 1.1 Ga (Kumar et al., 2007).
The age of the Majhgawan lamproite body (1067 ± 31 Ma) located 1000 km to the
north of the Wajrakarur kimberlite field, fuelled speculation that ultrapotassic,
alkaline and mafic magmatism at this time occurred on a greater scale across India.
Kumar et al. (2007) suggested that the 1.1 Ga potassic-ultrapotassic and alkaline
mafic-magmatism in India was a part of a global geodynamic event as kimberlites and
lamproites of similar ages are recognized worldwide in countries such as Australia,
Greenland, Liberia, North America, Scandinavia and South Africa.

This study presents robust new ages for the emplacement of some southern Indian
kimberlites and lamproites and uses Ar-Ar age dating techniques on phlogopite
separates. Our work focuses predominantly on the Krishna lamproites, which in
particular lack reliable data to constrain their emplacement age(s). These age data are
then combined with geochemical data from the kimberlites and lamproites to provide
further constraints on the source region, and primary magma composition, prior to
kimberlite and lamproite emplacement. Our new age data for the kimberlites and
lamproites also provides insight into the geodynamic evolution of this region.

2. The southern Indian kimberlites and lamproites

The majority of kimberlite and lamproite intrusions in the region occur within the
Eastern Dharwar craton (EDC) (Fig.1). Kimberlites crop out to the west of the
Cuddapah Basin, while lamproites intrude the basin and its north-eastern margin.
Among these are the Krishna lamproites (Fig. 1), which are thought to be some of the
oldest lamproite occurrences globally (Chalapathi Rao et al., 2004), and speculated to be the sources of several of the notable Indian diamonds (Chakrabarti et al., 2007).

The EDC is primarily composed of ancient (>2 Ga), greenschist- to granulite-facies schists and gneisses (Chalapathi Rao et al., 2004). The Eastern Ghat orogeny (1.3-1.6 Ga) affected the EDC, resulting in a narrow, highly deformed granulite-facies belt extending from Chennai to near Kolkata. The ~2.0 Ga old Cuddapah Basin (Anand et al., 2003) covers an area of around 44000 km$^2$ (Nagaraja Rao et al., 1987) and lies within the EDC. It comprises a 6 to 12 km thick succession of igneous and sedimentary rocks of Early to Late Proterozoic age.

The kimberlites in the EDC predominantly occur in two spatially separated groups: the diamondiferous Wajrakarur kimberlite field (WKF) and the Naryanpet kimberlite field (NKF) (Fig. 2).

The Cuddapah Lamproite Field (CLF) comprises two lamproites - Chelima and Zangamarajupalle. The 30 known lamproites from the Krishna Lamproite Field (KLF) (Fig. 3) occur mainly as dikes and are hosted by early Proterozoic granites of the Peninsular Gneissic Complex of the Dharwar Craton.

2.1 Previous age determinations of kimberlites and lamproites in southern India

All radiometric ages determined for kimberlites and lamproites from this region have yielded Proterozoic ages (see Table 1 and references therein); however, it is vigorously debated whether the southern Indian kimberlites are contemporaneous, or
whether they represent separate magmatic episodes (e.g. Chalapathi Rao et al., 2004; Kumar et al., 2007).

Kumar et al. (2007) argued for a contemporaneous emplacement at around 1.1 Ga for all the kimberlites in the Eastern Dharwar Craton. In contrast, Chalapathi Rao et al. (1996, 1999) used their age data (Table 1) to suggest that there were two episodes of Proterozoic mafic potassic magmatic activity in the EDC: one at ca. 1.4 Ga for the NKF and a second at 1.1 Ga for the WKF. However, further age data (Table 1) for the NKF using various techniques (Rb-Sr and U-Pb; Kumar et al. 2001, 2007; K-Ca; Gopalan and Kumar, 2008) indicated that the NKF and WKF were both emplaced at ~1100 Ma.

There are few ages for the Indian lamproites, especially for those from EDC (Table 1) and those that are published are considered unreliable because of alteration effects in the kimberlite and lamproites. Only one lamproite from the KLF (Ramannapeta) has been dated previously. Kumar et al. (2001) reported a Rb-Sr age for the Rammanapeta lamproite (1224 ± 14 Ma), which although contradicts the K-Ar age (1384 ± 18 Ma) reported by Chalapathi Rao et al. (1996), has been used more frequently to represent the emplacement age of the entire KLF (e.g. Chakrabarti et al., 2007). This is because the possibility that excess Ar in mantle-derived phlogopite causes the older K-Ar ages (as shown by some Siberian and South African kimberlites e.g., Pearson et al., 1995) cannot be ruled out.

For the Chelima lamproite, the ages of Chalapathi Rao et al. (1996) and Kumar et al. (2001) are in agreement. However, Kumar et al. (2001) suggested that their Rb-Sr ages
for Chelima and Zangamarajupalle were only tentative, as these samples display extensive secondary carbonation, which may have adversely affected the “true” crystallization ages of the lamproites. The K-Ar age reported by Chalapathi et al. (1996) has also been considered unreliable (Kumar et al., 2005) due to the possibility of excess Ar in mantle-derived phlogopite leading to the older K-Ar ages.

From Table 1 and the preceding discussion it is apparent that the recent age determinations provide a compelling argument for contemporaneous kimberlite emplacement in the EDC at around 1100 Ma. However, little reliable data exists for the lamproites from Southern India, and in particular the KLF; the published ages (Table 1) suggest that the kimberlites are not contemporaneous with the older lamproites.

In this study we employed the Ar-Ar method to determine new age data to better constrain the age of the KLF. Whilst the use of the K-Ar and Ar-Ar methods to date the kimberlites and lamproites of southern India has been questioned (Kumar et al., 2007), studies elsewhere have successfully dated kimberlites and lamproites through applying the Ar-Ar dating technique to phlogopite (Phillips 1991), yielding ages which were consistent with previous age determinations using different methods. Lehmann et al. (2010) recently used the Ar-Ar age dating technique on phlogopite separates as well as U-Pb on perovskites to date kimberlites in the MKF, with both methods providing a similar age. We have also obtained new dates on some kimberlites from the EDC which agree with well-constrained published ages (Chalapathi Rao et al., 1999, Kumar et al., 2007). Comparison of our data with
previously published ages allows us to assess the reliability of our age data for the KLF.

3. Samples, methodology and analytical techniques

3.1 Samples analysed and assessment of alteration

Samples were collected from the interior of each exposure (locations listed in Table 2). The kimberlites are classified as ‘Group I’ kimberlites on the basis of their mineralogy, in agreement with previous studies (e.g. Chalapathi Rao et al., 2004). Kimberlites and lamproites are susceptible to alteration processes and crustal contamination, and given the Proterozoic age of the rocks in this study, any interpretation of geochemical data must first consider the potential effect of these processes on the samples.

To assess any contamination or alteration effects, we first used the Ilmenite Index of Taylor et al. (1994), which identifies kimberlites and lamproites that may have accumulated or assimilated ilmenite megacrysts and xenocrysts. We also used loss on ignition (LOI) data as a proxy for alteration (Chalapathi Rao et al., 2004). Samples with high LOI are considered indicative of secondary alteration, commonly manifested in thin sections through increased abundances of secondary carbonate minerals and talc. Gd/Lu ratios of samples were also considered, where low values are indicative of HREE-enriched crustal contaminants (le Roex et al., 2003). A cut-off value of <58 was considered contaminated (Paton et al., 2009). The contamination index (CI, a measure of the proportions of clay minerals and tectosilicates relative to ferro-magnesian minerals - olivine, phlogopite) of Clement (1982) was used in this study, where a CI value of <1.5 is considered uncontaminated for kimberlites.
(Mitchell, 1986). The kimberlite samples used in this study are below this cut-off value; however the lamproites show higher values, which are not unusual for such mica-rich rocks (Chalapathi Rao et al., 2004). CI values in the whole suite of samples from the Krishna lamproites ranged from 3.79-9.23.

Care was taken to ensure that all the samples we collected from the EDC were as fresh as possible. Those chosen for dating in this study are among the freshest and showed minimal contamination and alteration as identified by CI, LOI, Gd/Lu and Ilmenite Index determinations for our sample suite (Table 2).

3.2 Ar-Ar dating technique

Age data for three samples are presented here, including two kimberlites (Muligiripalle Pipe 5 and Tummatapalle pipe 13) from the WKF and one lamproite (Pochampalle) from the KLF. The Pochampalle lamproite is a NW-SE trending body located 2.5 km west of the Pochampalle village. Petrographic analysis of thin sections identified samples with the most suitable phlogopite grains for dating. The samples with freshest phlogopite were selected, with several samples discounted on the basis of high calcite content and alteration of phlogopite to chlorite. Once selected, microprobe analysis of phlogopite within each selected sample was used to determine the mineral chemistry.

Small blocks of each of the selected samples were then sawn, avoiding any crustal xenoliths, and crushed in a jaw crusher. The fragments were then sieved into fractions from which the phlogopite could be carefully picked (predominantly 300μm-500μm in length). The picked phlogopite grains were then washed in acetone using an
ultrasonic bath to remove any adhering material. The visibly fresh grains with fewest inclusions were then selected using a binocular microscope. These grains were then packaged in aluminium foil and sent for sample irradiation at McMaster University in Ontario, Canada. Irradiation flux was monitored using the GA1550 biotite standard with an age of 98.79 ± 0.54 Ma (Renne et al., 1998). Sample J values were calculated by linear interpolation between two bracketing standards and are included in Table 3; a standard was included between every 8-10 samples in the irradiation tube. Blanks were measured either side of each measurement and used to correct each unknown, and \(^{37}\text{Ar}\) decay and neutron-induced interference reactions using the correction factors \((^{39}\text{Ar}/^{37}\text{Ar})_{\text{Ca}} = 0.00065 \pm 0.000033\), \((^{36}\text{Ar}/^{37}\text{Ar})_{\text{Ca}} = 0.000264 \pm 0.000013\) and
\((^{40}\text{Ar}/^{39}\text{Ar})_{\text{K}} = 0.0085\), and the mass discrimination value used was 283. The decay constant of Steiger & Jäger (1977) was used.

Samples were loaded into an ultra-high-vacuum laser port and placed under a heat lamp for 8 hours to reduce atmospheric blank levels. Samples were analysed by total fusion of single grains using Nd-YAG 1064 nm infrared laser, or Nd-YAG 213 nm UV laserprobe, both coupled to a MAP 215-50 mass spectrometer. Gases were gettered for 5 minutes using two SAES getters one at 450°C and one at room temperature, and a liquid nitrogen cold trap, before inlet to the mass spectrometer. Peaks between \(^{35}\text{Ar}\) and \(^{41}\text{Ar}\) were scanned 10 times and amounts extrapolated back to the inlet time. Each analysis was background corrected using blank measurements bracketing every 1-2 samples.

The infrared-laser single-grain-fusion technique was applied to Muligiripalle Pipe 5 (Mul) and Tummatapalle pipe 13 (Tum) and Pochampalle (POCg) samples. Here
individual mineral grains were fused to yield a single age, and the final age is a weighted mean average, error and MSWD as calculated using Isoplot 3a (Ludwig, 2003) and reported at 2σ level. The UV intragrain laserprobe technique was applied to POC and POCg samples, in which mineral grains were large enough to enable several measurements within each grain in order to test for heterogeneity (Figure 4.) (e.g., Kelley & Wartho, 2000; Sherlock et al., 2002). Each age determination was derived from a rastered laser pit measuring 75μm x 75μm, with a 10 μm diameter beam and reported at 2σ level. In addition, grain margins were analysed using trenches parallel to the grain margins with a beam size of 10μm. The number of analysis on each grain depended on its size, but can be seen in Figure 4. (The full dataset is available in online data repository where all values are reported in voltages)

3.3 Bulk-Rock and Isotopic measurements protocol

1-2 kg of visibly unaltered kimberlite and lamproite samples were cut on a rock saw to remove any weathered portions and then reduced to small fragments in a jaw-crusher. These fragments were then crushed to powder in an agate ring mill to produce the powders used in all bulk-rock geochemical analysis.

Two methods were used to obtain bulk-rock major-, trace- and REE data. Major and some minor elements (Al, Ca, Fe, K, Mg, Mn, Na, P, Si, Ti, Ba, Cr, Ni, Sr, Y, Zr), were analysed using Inductively-Coupled Plasma Atomic Emission Spectroscopy (ICP-AES). 100 mg of powdered sample was fused with 300 mg LiBO$_2$ (lithium metaborate) flux, and then dissolved in dilute HNO$_3$. The concentrations of selected major- and trace-elements were then determined by ICP-AES at the Natural History Museum, London. These solutions were not spiked. Reference materials were selected
based on their similarities to the samples to be analysed and included BCR-1 (major
and trace) and GA (trace and REE).

HF/HClO₄ (hydrofluoric acid/perchloric acid) digestion was used for other minor-,
trace- and REE analyses. In each case, 500 mg of powdered sample underwent a total
digestion in HF/HClO₄. The resulting solution was evaporated to dryness, and the
residue was then dissolved in dilute nitric acid. The concentrations of trace-elements
were then determined by ICP-AES and ICP-MS, as required, at the Natural History
Museum, London. Solutions for ICP-MS were diluted a further 10 times and were
spiked with 1 ppb In and 1 ppb Rh as internal standards prior to analysis. ICP-AES
solutions were not spiked. To monitor instrumental drift and precision the standard
BCR-1 was used. Major, trace and REE element analysis were found to be better than
± 20 % (2 s.d.) to that of the certified values, with the exception of Ni, Cr, Sn and Be
which were found to be better than ± 30 % (2 s.d.) to that of the certified values and
Bi, Cd, and Ta which showed slightly greater deviation to that of certified values.

The four dated samples underwent Nd isotope analysis. 50 mg of powdered sample
was weighed in each case and spiked with a ¹⁵⁰Nd/¹⁴⁴Nd solution. These samples
underwent a total digestion in HF/HNO₃. Nd and Sm were separated from the
dissolved samples by standard ion-exchange techniques (Cohen et al., 1988), using
cation columns to collect the REE fraction, which was then passed through HDEHP
columns to collect Nd and Sm fractions, which were evaporated to dryness. 1μl of 2M
HCl was added to the Nd collected for each sample and then loaded with 0.6μl of
0.01M H₃PO₄ onto the centre of an outgassed Re filament. Nd isotopic ratios were
measured in the static mode using a Triton thermal ionization mass spectrometer at
the Open University. Reproducibility of the La Jolla Nd standard over the analysis period was $0.511849 \pm 0.000002147$ (8 ppm, 2SD)

4. Results

4.1 Ar-Ar dating

The Ar-Ar dating results are summarised in Table 3 and reported to $2\sigma$. Where infrared laserprobe analysis is performed on mineral separates each age is derived from the total fusion of an individual mineral grain.

4.1.1 Kimberlites: Muligiripalle Pipe 5 and Tummatapalle Pipe 13

For the Muligiripalle Pipe these ages range from 1088±5 to 1149±21 Ma (n=18) with a weighted mean age of 1113±3 Ma (MSWD=0.96; n=17). For the Tummatapalle Pipe the age range from 1098±16 to 1138±29 Ma (n=10), with a weighted mean age of 1105±12 (MSWD=0.24; n=10).

4.1.2 Lamproite: Pochampalle

Infrared laserprobe analysis yields individual mineral ages in the range 1408±5 to 1614±8 Ma (n=6). With such a scatter in age data it is important to assess the spatial variation of ages within individual mineral grains which is achievable using the ultraviolet intra-grain technique. Five different mineral grains from two samples have been analysed using this technique (Figure 4). From sample POC ages within the internal parts of the mineral grains range from 1543±39 to 1611±32 Ma, and from the grain margins range from 1480±35 to 1625±15 Ma (n=8). From sample POCg ages within the internal parts of the mineral grains range from 1463±71 to 1684±100 Ma (n=24) and from the grain margins range from 1507±38 Ma to 1698±30 Ma (n=11).
4.2 Geochemistry

4.2.1 Geochemical and isotopic data on the Krishna lamproites

Major, trace and REE data were obtained on lamproites from 9 different locations (the full dataset is available in the online data repository). Two samples were analysed from two of the lamproites, Vedadri (VEDN and VEDS) and Pochampalle (POC and POCg); in each case, the two samples were collected from different locations within the same lamproite body. 8 of the lamproite samples are from the KLF, while one (Chelima) is located within the Cuddapah Basin (Table 2).

All of the lamproites in this study are LREE enriched, as shown by elevated ($\text{La/Asb}_N$) values (Figure 5). For the Krishna lamproites, ($\text{La/Asb}_N$) = ~ 45 to 75. The Pochampalle lamproite shows even greater LREE enrichment compared to the rest of the KLF samples with ($\text{La/Asb}_N$) = ~100. The Chelima lamproite has the highest degree of LREE enrichment among the sample suite analyzed in the present study, with ($\text{La/Asb}_N$) = ~200. These values are comparable to other published ratios for the KLF ($\text{La/Asb}_N$) = 51-173 (Chalapathi Rao et al., 2010) and Chelima ($\text{La/Asb}_N$) = 72-247 (Chalapathi Rao et al., 2004).

On a chondrite-normalized REE plot (Figure 6), all lamproites from the EDC display similar LREE-enriched patterns. The Chelima lamproite shows the steepest REE pattern. Two samples were collected from different locations from the Pochampalle lamproite and show slightly different REE patterns (Fig 6). The POC sample shows a trend closer to those of the KLF, especially for the HREE, though with a higher La/Yb ratio due to greater LREE enrichment. The POCg sample has an even higher La/Yb
ratio due to its extreme LREE enrichment (similar to that of Chelima), but has higher HREE contents than Chelima. The HREE enrichment may be due to variations in the source but can also be attributed to crustal contamination. However, even if the samples have undergone some degree of crustal contamination, it is unlikely to have affected the isotopic ratios of small-fraction melts, because the concentrations of Sm and Nd in crustal rocks is likely to be much lower than in the kimberlitic melts (e.g. Fraser et al., 1985; Gibson et al., 1996).

The Pochampalle ($^{143}\text{Nd}/^{144}\text{Nd} = 0.511059$) and Chelima ($^{143}\text{Nd}/^{144}\text{Nd} = 0.511129$) lamproites also show lower Nd isotopic values than the rest of the lamproites in the KLF ($^{143}\text{Nd}/^{144}\text{Nd} = 0.511348-0.511459$) (Figure 7). In contrast to the REE data, the $^{143}\text{Nd}/^{144}\text{Nd}$ isotopic ratio of Pochampalle is not intermediate between that of Chelima and the Krishna lamproite group.

The lower $^{143}\text{Nd}/^{144}\text{Nd}$ ratio of the Pochampalle lamproite suggests a closer affinity with the CLF, particularly Chelima, than with the rest of the KLF samples. The broad spread of the KLF Nd data precludes robust interpretations of a mixing array involving the Cuddapah Basin Lamproites (CBL), Pochampalle and the Krishna lamproites. However, the difference in Nd isotopic signature between Pochampalle and the other Krishna lamproites is striking.

The Nd evolution through time of the southern Indian kimberlites and lamproites in this study is summarised in Figure 8, and compared with model evolution curves for depleted mantle (DM; Workman and Hart, 2005), and enriched mantle (EM). The enriched mantle is modelled using the $\varepsilon\text{Nd}$ value of a garnet lherzolite xenolith from
the Lattavaram kimberlite in the EDC (Karmalkar et al. 2009), and assuming a representative value of $^{143}$Nd/$^{144}$Nd for enriched mantle (EMI; Hart et al., 1992).

The $\varepsilon_{Nd}$ evolution trajectory of the kimberlites intersects the EM* model evolution line at around 1 Ga, which corresponds to the emplacement age of the EDC kimberlites. However, the DM and EM* model evolution lines intersect much earlier (1.9-2.0 Ga). One model for petrogenesis of the EDC kimberlites consistent with these Nd evolution data is that the kimberlite magmas were derived at around 1 Ga from an initially depleted mantle source that had been enriched at around 1.9-2 Ga.

The mantle source from which the lamproites were derived is not tightly constrained by the evolution curves in Figure 8. Their enriched geochemistry indicates that they did not originate directly from a depleted mantle source, but clearly they have a different source to that of the kimberlites. It is also apparent that the Pochampalle and Chelima lamproites followed a separate evolution from the other Krishna lamproites.

It is possible that Pochampalle and Chelima lamproites were derived from a different source altogether from other Krishna lamproites. However, they may have been derived from the same source as the Krishna lamproites but at an earlier time. This source could have been an ancient depleted mantle that was enriched prior to lamproite generation. More definite conclusions are difficult to draw from Nd evolution modelling of the lamproites due to the isotopic heterogeneity of the sub-continental lithospheric mantle, even on relatively small length scales.

5. Discussion
Despite the recent increase in scientific interest in the kimberlites and lamproites of the Dharwar Craton in southern India, the ages of the kimberlites and in particular lamproites are poorly constrained. An age of around 1.1 Ga for both the kimberlites in the WKF and NKF appears to be well supported, although many pipes within these fields are yet to be dated. The only lamproite previously dated from the KLF is the Ramannapeta lamproite, even though there are many lamproites in the KLF; and it would seem presumptuous to assume an age for the entire field based on one age from one lamproite. However, this age has been used in several studies to represent an age for the KLF mainly because of a lack of any other reliable age data (Reddy et al., 2003, Paul et al., 2007, Chakrabarti et al., 2007).

The Ar-Ar ages for Muligiripalle of 1113 ± 3 Ma and for Tummatapalle of 1105 ± 12 are consistent with previous age determinations for the WKF and in particular Muligiripalle, which has been dated at 1093 ± 20 (Kumar et al., 1993) by Rb-Sr technique and 1153 ± 17 (Chalapathi Rao., 1996) by K-Ar technique. The Pochampalle lamproite from the KLF yields a much older age than the published K-Ar and Rb-Sr age dates for the KLF (1224 ± 14 Ma on the Ramannapeta lamproite; Kumar et al., 2001). However, both the K-Ar and Rb-Sr ages were described as tentative by the authors as they were unable to rule out the effects of excess extensive secondary carbonation (Rb-Sr) and the possibility of excess Ar in mantle-derived phlogopite (K-Ar). We dated samples from two different locations from within the Pochampalle lamproite (POCg and POC). The dating of kimberlites and lamproites has been the subject of discussion, because in many cases phlogopite cores are older than their rims, which is true of the Pochampalle lamproite. In rocks formed and cooled in mid- and upper-crustal settings this is usually attributed to either prolonged
cooling imparting an ‘age’ gradient or partial resetting of a mineral and argon-loss occurring at the rims (e.g. Sherlock et al., 2002). In kimberlites and lamproites the same observed age patterns are argued to be due to a pervasive and uniformly-distributed excess argon component derived from a deep fluid source (e.g. Phillips & Onstott, 1988; Phillips 1991; Phillips et al., 1999) that is then partially outgassed to the grain boundary network during incorporation into the magma chamber (Pearson et al., 1997). Phillips & Onstott (1988) compared step-heating spectra of two sets of phlogopite mineral separates with \textit{in situ} data obtained from a single large phlogopite grain using a continuous wave infrared Nd-YAG laser. The step-heating spectra identify the range of ages preserved within the mineral separates but without identifying the spatial distribution of the ages, whilst the laserprobe data identifies the position of the ages within the grain. The resolution of the laserprobe data is limited by the area outside the point of laser-sample interaction that is outgassed due to heat conduction during lasing; Phillips & Onstott (1988) report a resolution of 80-150\textmu m diameter pits separated by distances of 20-100\textmu m, which gave sufficient confidence that the individual age measurements were not affected by adjacent laser pits. The age range of ca. 1200 to ca. 1540 Ma from low- to high-temperature steps in the heating spectra indicate a range of ca. 1200 Ma to ca. 2200 Ma (Phillips & Onstott, 1988), the oldest age represented by a single laser spot. The differences were difficult to reconcile but it was considered that the old ages could be ‘masked’ during physical mineral decrepitation during step-heating and the mixing of different aged reservoirs within the mineral separates. Notably, the volume of phlogopite preserving old apparent ages is a small fraction of the grain and so it is conceivable that during the preparation of the mineral separates by crushing and sieving this low-volume component might not be well represented. The study raised more questions about the
behaviour of argon in such high pressure and high temperature minerals. Phillips (1991) also compared step-heating and laserprobe Ar-Ar data, and analysed matrix and macrocryst phlogopites. The former yielded plateau ages that were consistent with eruption ages and the latter yielded complex release patterns concurring with those reported by Phillips & Onstott (1988). In contrast the laserprobe data (spatial resolution of 100x200µm sized pits in the internal parts of the grain and up to 160x400µm at the rim, separated by 30-80µm) reveal older cores and younger rims, with similar conclusions to Phillips & Onstott (1988), in that a high concentration of excess argon was trapped in the macrocrysts prior to eruption, the rims record the eruption age, and the matrix phlogopites record the eruption age (Phillips, 1991). This model dictates that radiogenic argon is retained in minerals in the upper mantle at temperatures of 700°C or more (Pearson et al., 1997). Kelley & Wartho (2000) were able to test this by analysing well-characterised xenoliths from two different settings, with robust age constraints, using the high spatial resolution ultra-violet laserprobe approach to assess core-rim age differences in detail. The results showed old cores and young rims, and in both settings the older core ages corresponded to magmatic/metasomatic events that were recorded by other isotope systems (Kelley & Wartho, 2000). The rims were also younger than the cores, with argon-loss profiles of 200 to 300µm in length decreasing to the known eruption age at the grain edge (Kelley & Wartho, 2000). This study identified that the phlogopites retained argon at temperatures hundreds of degrees higher than the phlogopite closure temperature (ca. 450° C) and that argon loss at the grain margins represents the integrated time-temperature history as the xenolith travels from depth to emplacement at the surface (Kelley & Wartho, 2000). The key difference between this study and the conclusions of Phillips et al (1991; 1999) is in the interpretation of the core-rim age differences:
Phillips (1991) and Phillips et al (1999) interpret old cores due to excess argon whilst Kelley & Wartho (2000) interpret them as retained radiogenic argon because of a lack of transfer of argon to the grain boundary network because the grains behave as ‘closed systems’ under the high pressures experienced in the mantle and lower crust.

In terms of our new data for the Pochampalle lamproite, macrocrystic phlogopite yielded single grain fusion ages that spanned ca. 200 Myr, and the intra-grain ultraviolet laserprobe analysis revealed a complex age structure with older internal parts and varied rim ages. Individually the five grains are different, in size and shape, but the age ranges for each of the grains are very similar, most notably for sample POC. Contouring is not appropriate in this case because there are two analytical approaches – ablating squares in internal parts of the grains and margin-parallel trenches – and taking the approach of plotting age versus distance from the centre of the laser pit to the nearest grain margin can reveal information on the age structure (Figure 9).

Sample POC preserves cores of ca 1560 Ma to ca 1610 Ma with a single data point that is much higher (ca. 1660 Ma). In both grains the measurements actually at the grain margins are the youngest – ca. 1500 Ma and ca. 1480 Ma. In POCg the same age pattern is observed though with more scatter, and the key point is that in plotting data in this way it assumes that the distance from laser pit to grain margin now, is the same as it was at the time of mineral growth, a likely false assumption given the difficulty in recovering intact mineral grains. Notably for POCg the grain margins are consistently the youngest with ages of ca 1493 Ma, ca. 1463 Ma and ca. 1507 Ma, although there is notable scatter in the third grain from sample POCg in which two grain margins are measured, one of which is significantly older than the other and
may not represent an original grain margin. The internal parts of the grain are consistently in the range ca. 1530 Ma to ca. 1630 Ma. The key points to note are: 1) in five mineral grains from samples POC and POCg the grain margins are the youngest parts of the grains and are all within error of 1500 Ma; 2) the internal parts of the mineral grains are significantly older and in the region of ca. 1550-1650 Ma. These observations are in keeping with those of Phillips & Onstott (1988), Phillips (1991), Phillips et al (1999) and Kelley & Wartho (2000). Whether the older core regions of the minerals reflect excess argon or retention of radiogenic argon under mantle-lower crust conditions is difficult to discern, but by and large this is a moot point. It is the ages recorded by the rims of the minerals that are of importance since these are considered to represent the timing of eruption by all the previously cited studies.

To summarise the new Ar-Ar data, ages from the two kimberlite pipes derived from matrix phlogopites are consistent and yield an age of ca. 1100 Ma for their eruption. The Pochampalle lamproite preserves old ages in internal parts of the grain that could be due to either excess argon retention or quantitative retention of radiogenic argon, whilst the rims record an eruption age of ca. 1500 Ma.

Based on these Ar-Ar age data, we propose that the Pochampalle lamproite was emplaced ~1500 Ma ago, roughly 250 Myr before the rest of the KLF. It would seem that the Pochampalle magma was also derived from an isotopically distinct mantle source from the rest of the KLF lamproites with a lower $^{143}\text{Nd}/^{144}\text{Nd}$ ratio. Trace element geochemistry implies that it may have been derived by relatively smaller degrees of partial melting from within the garnet stability field compared to the rest of the KLF. The Nd isotopic signature and the bulk-rock REE pattern in the Pochampalle
lamproite appear to be consistent with a hypothesis that a heterogeneous mantle source was sampled by the Krishna lamproites during the Proterozoic.

Insight into the possible tectonic setting for emplacement of the Pochampalle lamproite could come from recent work by Hou et al. (2008). Citing a giant radiating dyke swarm and LIPs at ~1.85 Ga, they propose that the North China Craton, Indian Craton and Canadian Shield were united together in a single landmass before its extension and break-up. The Mesoproterozoic Belt–Purcell–Uinta rift system (1470–1440 Ma) (Sears et al., 1998) along the west margin of the Canadian Shield suggests that the North China Craton and the Indian Craton were ultimately separated from the Laurentia continent by around 1.5 Ga onwards. This is close to the emplacement age of the Pochampalle lamproite, which could therefore be linked to this period of continental rifting.

The older age and smaller degrees of mantle partial melting inferred for the Pochampalle lamproite may correspond to initiation of lithospheric stretching beneath the southern Indian craton at ~1500 Ma. The continuation or pre-existence of such a weakness in the lithosphere may have been exploited by later emplacement of the Krishna lamproites, by which time the mantle source region had acquired modified isotopic signatures through melt percolation from the asthenospheric mantle.

Previously, the Chelima lamproite (ca. 1400 Ma) was thought to be one of the oldest recorded lamproites in the world (Chalapathi Rao, 2007). However, our age data suggest that at least one lamproite (Pochampalle) was generated in the same region 100 Ma before the Chelima lamproite. This not only has implications for regional
ultramafic magmatism, but also demonstrates that the mantle mechanism for producing lamproitic melts existed earlier than previously thought. It seems likely that further age determinations on Indian lamproites may extend their age range even further.

Acknowledgements

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References


Figure Captions

Figure 1. Location of the Krishna lamproites relative to the major tectonic domains of southern India. HKF: Hinota Kimberlite Field; MKF: Mainpur Kimberlite Field; NKF: Naryanpet Kimberlite Field; WKF: Wajrakarur Kimberlite Field. Modified from Paton et al. (2009).

Figure 2. Location and extent of kimberlite and lamproite fields around the Cuddapah Basin, showing the locations of individual pipes. Modified from Chalapathi Rao et al. (2009). The inset shows the location of the KLF expanding north to provide a more detailed map of the Krishna lamproite field (KLF) showing the location of the individual lamproite pipes including Pochampalle (KLP on diagram). Modified from Chakrabarti et al. (2007).

Figure 3. Typical microtextures of phlogopite in the analysed samples. Each set shows a back-scattered electron image (left) and an element map for potassium (right). a) Muligiripalle kimberlite (pipe 5, WKF), scale bars 200 μm; b) Tummatapalle kimberlite (pipe 13, WKF), scale bars 100 μm; c) Pochampalle lamproite (KLF), scale bars 500 μm.

Figure 4. Sketches of phlogopite grains analysed using UV intragrain laserprobe technique. Each age determination was derived from a rastered laser pit measuring 75 μm x 75 μm, with a 10 μm diameter beam. The number of analyses on each grain depended on its size.

Figure 5. Plot of Dy/Yb against La/Yb (normalized to chondrite; values from Anders and Grevesse, 1988) for the Krishna lamproites and Chelima lamproite. Data from Chak (Chakrabarti et al., 2007) and Chal (Chalapathi Rao et al., 2004) are also shown.

Figure 6. Chondrite-normalized (values from Anders and Grevesse, 1988) REE plot for the Krishna lamproites (average abundance), the Chelima lamproite (solid line), and the two Pochampalle lamproite samples (dashed lines).

Figure 7. Plot of 1/[Nd] against 143Nd/144Nd ratios for the Krishna lamproites (including Pochampalle) and the CLF (including Chelima). Data are from this study, Chakrabarti et al. (2007) (Chak) and Chalapathi Rao et al. (2004) (Chal).

Figure 8. Plot of εNd against time for the lamproites (solid lines) from the EDC (data from this study). The Pochampalle and Chelima lamproites are shown as bold solid lines. Also shown is the range of kimberlites data from the EDC (shaded field; Paton et al., 2009). The single bold dashed line represents the potential Nd evolution of enriched mantle (EM*), beneath India, modelled using the εNd of a garnet lherzolite xenolith from the Lattavaram kimberlite in the EDC (Karmalkar et al. 2009), combined with a representative value of 143Nd/144Nd for EMI (Hart et al., 1992). The depleted mantle (DM) trend (dotted line) uses data from Workman and Hart (2005).

Figure 9. Age versus distance from grain edge for all UV laserprobe data points from phlogopite macrocrysts from: a) Sample POC; b) Sample POCg.
**Table 1.** Compilation of dates obtained on kimberlites and lamproites from the Eastern Dharwar Craton and related areas

<table>
<thead>
<tr>
<th>Date (Ma)</th>
<th>Method</th>
<th>Material</th>
<th>Samples</th>
<th>Field</th>
<th>Authors</th>
</tr>
</thead>
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<td></td>
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</tr>
<tr>
<td>1093 ±20</td>
<td>Rb-Sr</td>
<td>phlogopite Muligiripalle pipe 5</td>
<td>WKF</td>
<td>K et al. 93</td>
<td></td>
</tr>
<tr>
<td>1153 ±17</td>
<td>K-Ar</td>
<td>phlogopite Muligiripalle pipe 5</td>
<td>WKF</td>
<td>CR et al. 96</td>
<td></td>
</tr>
<tr>
<td>1091 ±20</td>
<td>Rb-Sr</td>
<td>phlogopite Wajrakarur pipe 1</td>
<td>WKF</td>
<td>K et al. 93</td>
<td></td>
</tr>
<tr>
<td>1092 ±15</td>
<td>Rb-Sr</td>
<td>phlogopite Wajrakarur pipe 2</td>
<td>WKF</td>
<td>K et al. 93</td>
<td></td>
</tr>
<tr>
<td>1102 ±23</td>
<td>Rb-Sr</td>
<td>phlogopite Wajrakur</td>
<td>WKF</td>
<td>K et al. 07</td>
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<tr>
<td>1124 ±5</td>
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<td>WKF</td>
<td>K et al. 07</td>
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<td>Rb-Sr</td>
<td>phlogopite Venkatampalle pipe 7</td>
<td>WKF</td>
<td>K et al. 93</td>
<td></td>
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<td><strong>Narayanpet Kimberlite Field (NKF)</strong></td>
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<td></td>
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<td></td>
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<tr>
<td>1363 ±48</td>
<td>K-Ar</td>
<td>phlogopite Kotakonda</td>
<td>NKF</td>
<td>CR et al. 96</td>
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<tr>
<td>1401 ±5</td>
<td>Ar-Ar</td>
<td>phlogopite Kotakonda</td>
<td>NKF</td>
<td>CR et al. 99</td>
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<td>1085 ±14</td>
<td>Rb-Sr</td>
<td>phlogopite Kotakonda</td>
<td>NKF</td>
<td>K et al. 01</td>
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<tr>
<td>1099 ±12</td>
<td>Rb-Sr</td>
<td>phlogopite Mudalabad</td>
<td>NKF</td>
<td>K et al. 01</td>
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</tr>
<tr>
<td>1167 ±86</td>
<td>K-Ca</td>
<td>phlogopite Narayanpet</td>
<td>NKF</td>
<td>G &amp; K 08</td>
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<tr>
<td>1093 ±4</td>
<td>Rb-Sr</td>
<td>phlogopite Siddanpalle</td>
<td>NKF (SKF)</td>
<td>K et al. 07</td>
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<tr>
<td>1063 ±41</td>
<td>K-Ca</td>
<td>phlogopite Siddanpalle</td>
<td>NKF (SKF)</td>
<td>G &amp; K 08</td>
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<td><strong>Mainpur Kimberlite Field (MKF)</strong></td>
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<td>60 – 65</td>
<td>Ar-Ar</td>
<td>Phlogopite &amp; Perovskite</td>
<td>Behradih Kodomali</td>
<td>MKF (~700 km N)</td>
<td>L et al. 2010</td>
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<td>phlogopite Majhgawan</td>
<td>HKF (1000 km N)</td>
<td>K et al. 93</td>
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<td><strong>Cuddapah Basin Lamproites (CB)</strong></td>
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<td>1418 ±8</td>
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<td>Sample</td>
<td>Isotope</td>
<td>Mineral</td>
<td>Location</td>
<td>Notes</td>
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<td>1354 ±17</td>
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<td><em>Chelima</em></td>
<td>CB</td>
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<td>1090</td>
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**Krishna Lamproite Field (KLF)**

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<th>Sample</th>
<th>Isotope</th>
<th>Mineral</th>
<th>Location</th>
<th>Notes</th>
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<td>phlogopite</td>
<td><em>Ramannapeta</em></td>
<td>KLF</td>
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<tr>
<td>1224 ±14</td>
<td>Rb-Sr</td>
<td>phlogopite</td>
<td><em>Ramannapeta</em></td>
<td>KLF</td>
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</table>

1 Kimberlites: plain type; lamproites: *italics*.
2 K et al. 93: Kumar et al. (1993); CR et al. 96: Chalapathi Rao et al. (1996); CR et al. 99: Chalapathi Rao et al. (1999); K et al. 01: Kumar et al. (2001); K et al. 07: Kumar et al. (2007); G & K 08: Gopalan and Kumar (2008); L et al., 2010: Lehmann et al. (2010).
Table 2. Location and Contamination/Alteration of Samples.

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<th>Name</th>
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<th>Latitude</th>
<th>CI</th>
<th>LOI % Wt Loss</th>
<th>Ilmenite Index</th>
<th>Gd/Lu</th>
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<tr>
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<td>77° 32' 06''</td>
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<td>6.24</td>
<td>0.59</td>
<td>74.50</td>
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<td>Tummatapalle (K)</td>
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<td>77° 30' 09''</td>
<td>1.19</td>
<td>8.62</td>
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<td>Pochampalle (L)</td>
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<td>80° 16' 61''</td>
<td>5.09</td>
<td>3.91</td>
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Table 3. Ar-Ar results.

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<th>Sample</th>
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<th>Method¹</th>
<th>Wtd Mean (Ma)</th>
<th>Error (±)</th>
<th>MSWD</th>
<th>Number of analyses</th>
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<td>1115</td>
<td>4</td>
<td>0.78</td>
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<td>Tummatapalle</td>
<td>Kimberlite</td>
<td>TF</td>
<td>1105</td>
<td>12</td>
<td>0.24</td>
<td>10 grains fused</td>
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<td>Lamproite</td>
<td>UV</td>
<td>1575</td>
<td>20</td>
<td>1.3</td>
<td>14 points on single grain (Fig. 4a)</td>
</tr>
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<td>Lamproite</td>
<td>UV</td>
<td>1568</td>
<td>55</td>
<td>11.3</td>
<td>6 points on single grain (Fig. 4b)</td>
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<td>Pochampalle</td>
<td>Lamproite</td>
<td>TF</td>
<td>1523</td>
<td>110</td>
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<td>6 grains fused</td>
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<td>Lamproite</td>
<td>UV</td>
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<td>25</td>
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<td>12 points on single grain (Fig. 4c)</td>
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<td>1583</td>
<td>16</td>
<td>2.5</td>
<td>15 points on single grain (Fig. 4e)</td>
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</tbody>
</table>

¹ TF = Total fusion analysis; UV = Ultraviolet laser in situ analysis
Figure

Fig. 4 POC analysis

Fig. 4a POC analysis

Fig. 4b POCg analysis

Fig. 4c POCg analysis

Fig. 4d POCg analysis
Figure

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